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(57) Abstract

Novel recombinant allergens are disclosed. The allergens are non-naturally occurring mutants derived from naturally-occurring allergens. The overall a-carbon backbone tertiary structure is essentially preserved. Also disclosed are methods for preparing such recombinant allergens as well as use of thereof.

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MUTANT RECOMBINANT ALLERGENS

FIELD OF THE INVENTION

The present invention relates to novel recombinant allergens, which are non-naturally occurring mutants derived from naturally occurring allergens. Further, the invention relates to a method of preparing such recombinant allergens as well as to pharmaceutical to compositions, including vaccines, comprising the recombinant allergens. In further embodiments, the present invention relates to methods of generating immune responses in a subject, vaccination or treatment of a subject as well as processes for preparing the compositions of the invention.

BACKGROUND OF THE INVENTION

Genetically predisposed individuals become sensitised (allergic) to antigens originating from a variety of environmental sources, to the allergens of which the individuals are exposed. The allergic reaction occurs when a previously sensitised individual is re-exposed to the same or a homologous allergen. Allergic responses 25 range from hay fever, rhinoconductivitis, rhinitis and asthma to systemic anaphylaxis and death in response to e.g. bee or hornet sting or insect bite. The reaction is immediate and can be caused by a variety of atopic allergens such as compounds originating from grasses, trees, weeds, insects, food, drugs, chemicals and perfumes.

However, the responses do not occur when an individual is exposed to an allergen for the first time. The initial 35 adaptive response takes time and does usually not cause any symptoms. But when antibodies and T cells capable of reacting with the allergen have been produced, any subsequent exposure may provoke symptoms. Thus, allergic responses demonstrate that the immune response itself can cause significant pathological states, which may be life threatening.

The antibodies involved in atopic allergy belong primarily to immunoglobulins of the IgE class. IgE binds to specific receptors on the surface of mast cells and basophils. Following complex formation of a specific allergen with IgE bound to mast cells, receptor cross-linking on the cell surface results in signalling through the receptors and the physiological response of the target cells. Degranulation results in the release of i.a. histamine, heparin, a chemotactic factor for eosinophilic leukocytes, leukotrienes C4, D4 and E4, which cause prolonged constriction of the bronchial smooth muscle cells. The resulting effects may be systemic or local in nature.

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The antibody-mediated hypersensitivity reactions can be divided into four classes, namely type I, type II, type III and type IV. Type I allergic reactions is the classic immediate hypersensitivity reaction occurring within seconds or minutes following antigen exposure. These symptoms are mediated by allergen specific IgE.

Commonly, allergic reactions are observed as a response to protein allergens present e.g. in pollens, house dust mites, animal hair and dandruff, venoms, and food products.

In order to reduce or eliminate allergic reactions, carefully controlled and repeated administration of allergy vaccines is commonly used. Allergy vaccination is traditionally performed by parenteral, intranasal, or

sublingual administration in increasing doses over a fairly long period of time, and results in desensitisation of the patient. The exact immunological mechanism is not known, but induced differences in the 5 phenotype of allergen specific T cells is thought to be of particular importance.

Antibody-binding epitopes (B-cell epitopes)

10 X-ray crystallographic analyses of Fab-antigen complexes has increased the understanding of antibody-binding epitopes. According to this type of analysis antibodybinding epitopes can be defined as a section of the surface of the antigen comprising atoms from 15-25 amino is acid residues, which are within a distance from the atoms of the antibody enabling direct interaction. The affinity of the antigen-antibody interaction can not be predicted from the enthalpy contributed by van der Waals interactions, hydrogen bonds or ionic bonds, alone. The 20 entropy associated with the almost complete expulsion of water molecules from the interface represent an energy contribution similar in size. This means that perfect fit between the contours of the interacting molecules is a principal factor underlying antigen-antibody high 25 affinity interactions.

Allergy vaccination

The concept of vaccination is based on two fundamental observations of the immune system, namely specificity and memory. Vaccination will prime the immune system of the recipient, and upon repeated exposure to similar proteins the immune system will be in a position to respond more rigorously to the challenge of for example a microbial infection. Vaccines are mixtures of proteins intended to be used in vaccination for the purpose of

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generating such a protective immune response in the recipient. The protection will comprise only components present in the vaccine and homologous antigens.

- S Compared to other types of vaccination allergy vaccination is complicated by the existence of an ongoing immune response in allergic patients. This immune response is characterised by the presence of allergen specific IgE mediating the release of allergic symptoms upon exposure to allergens. Thus, allergy vaccination using allergens from natural sources has an inherent risk of side effects being in the utmost consequence life threatening to the patient.
- Approaches to circumvent this problem may be divided in three categories. In practise measures from more than one category are often combined. First category of measures includes the administration of several small doses over prolonged time to reach a substantial accumulated dose. category of measures includes physical Second modification of the allergens by incorporation of the aluminium allergens into σel substances such as hydroxide. Aluminium hydroxide formulation has adjuvant effect and a depot effect of slow allergen release reducing the tissue concentration of active allergen components. Third category of measures include chemical modification of the allergens for the purpose of reducing allergenicity, i.e. IgE binding.
- The detailed mechanism behind successful allergy vaccination remains controversial. It is, however, agreed that T cells play a key role in the overall regulation of immune responses. According to current consensus the relation between two extremes of T cell phenotypes, Th1 and Th2, determine the allergic status of an individual. Upon stimulation with allergen Th1 cells secrete

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interleukines dominated by interferon-y leading to protective immunity and the individual is healthy. Th2 cells on the other hand secrete predominantly interleukin 4 and 5 leading to IgE synthesis and eosinophilia and the 5 individual is allergic. In vitro studies have indicated the possibility of altering the responses of allergen specific T cells by challenge with allergen derived peptides containing relevant T cell epitopes. Current approaches to new allergy vaccines are therefore largely 10 based on addressing the T cells, the aim being to silence the T cells (anergy induction) or to shift the response from the Th2 phenotype to the Th1 phenotype.

In WO 97/30150 (ref. 1), a population of protein molecules is claimed, which protein molecules have a distribution of specific mutations in the amino acid sequence as compared to a parent protein. From the description, it appears that the invention is concerned with producing analogues which are modified as compared 20 to the parent protein, but which are taken up, digested and presented to T cells in the same manner as the parent protein (naturally occurring allergens). Thereby, a modified T cell response is obtained. Libraries of modified proteins are prepared using a technique denoted PM (Parsimonious Mutagenesis). 25

In WO 92/02621 (ref. 2), recombinant DNA molecules are described, which molecules comprise a DNA coding for a polypeptide having at least one epitope of an allergen of 30 trees of the order Fagales, the allergen being selected from Aln g 1, Cor a 1 and Bet v 1. The recombinant molecules described herein do all have an amino acid sequence or part of an amino acid sequence that corresponds to the sequence of a naturally occurring allergen.

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WO 90/11293 (ref. 3) relates i.a. to isolated allergenic peptides of ragweed pollen and to modified ragweed pollen peptides. The peptides disclosed therein have an amino acid sequence corresponding either to the sequence of the 5 naturally occurring allergen or to naturally occurring isoforms thereof.

Chemical modification of allergens

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10 Several approaches to chemical modification of allergens have been taken. Approaches of the early seventies include chemical coupling of allergens to polymers, and chemical cross-linking of allergens using formaldehyde, etc., producing the so-called 'allergoids'. The rationale 15 behind these approaches was random destruction of IgE binding epitopes by attachment of the chemical ligand while retaining reducing IgE-binding immunogenicity by the increased molecular weight of the Inherent disadvantages of 'allergoid' complexes. production are linked to difficulties in controlling the process of chemical cross-linking and difficulties in analysis and standardisation of the resulting high molecular weight complexes. 'Allergoids' are currently in clinical use and due to the random destruction of IgE 25 binding epitopes higher doses can be administered as compared to conventional vaccines, but the safety and efficacy parameters are not improved over use of conventional vaccines.

More recent approaches to chemical modification of allergens aim at a total disruption of the tertiary structure of the allergen thus eliminating IgE binding assuming that the essential therapeutic target is the allergen specific T cell. Such vaccines contain allergen sequence derived synthetic peptides representing minimal T cells epitopes, longer peptides representing linked T

cells epitopes, longer allergen sequence derived synthetic peptides representing regions of immunodominant T cell epitopes, or allergen molecules cut in two halves by recombinant technique. Another approach based on this rationale has been the proposal of the use of "low IgE binding" recombinant isoforms. In recent years it has become clear that natural allergens are heterogeneous containing isoallergens and variants having up to approximately 25% of their amino acids substituted. Some recombinant isoallergens have been found to be less efficient in IgE binding possibly due to irreversible denaturation and hence total disruption of tertiary structure.

In vitro mutagenesis and allergy vaccination

Attempts to reduce allergenicity by in vitro site directed mutagenesis have been performed using several allergens including Der f 2 (Takai et al, ref. 4), Der p 20 2 (Smith et al, ref. 5), a 39 kDa Dermatophagoides farinae allergen (Aki et al, ref. 6), bee venom phospholipase A2 (Förster et al, ref. 7), Ara h 1 (Burks et al, ref. 8), Ara h 2 (Stanley et al, ref. 9), Bet v 1 (Ferreira et al, ref. 10 and 11), birch profilin 25 (Wiedemann et al, ref. 12), and Ory s 1 (Alvarez et al, ref. 13).

The rationale behind these approaches, again, is addressing allergen specific T cells while at the same time reducing the risk of IgE mediated side effects by reduction or elimination of IgE binding by disruption of the tertiary structure of the recombinant mutant allergen. The rationale behind these approaches does not include the concept of dominant IgE binding epitopes and it does not include the concept of initiating a new protective immune response which also involves B-cells

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and antibody generation.

The article by Ferreira et al (ref. 11) describes the use of site directed mutagenesis for the purpose of reducing 5 IgE binding. Although the three-dimensional structure of Bet v 1 is mentioned in the article the authors do not use the structure for prediction of surface exposed amino acid residues for mutation, half of which have a low degree of solvent exposure. Rather they use a method developed for prediction of functional residues in proteins different from the concept of structure based identification of conserved surface areas described here. Although the authors do discuss conservation of a-carbon backbone tertiary structure this concept is not a part of the therapeutic strategy but merely included to assess in vitro IgE binding. Furthermore, the evidence presented is not adequate since normalisation of CD-spectra prevents the evaluation of denaturation of a proportion of the sample, which is a common problem. The therapeutic strategy described aim at inducing tolerance in allergen specific T cells and initiation of a new immune response is not mentioned.

The article by Wiedemann et al. (ref. 12) describes the
use of site directed mutagenesis and peptide synthesis
for the purpose of monoclonal antibody epitope
characterisation. The authors have knowledge of the
tertiary structure of the antigen and they use this
knowledge to select a surface exposed amino acid for
mutation. The algorithm used can be said to be opposite
to the one described by the present inventors since an
amino acid differing from homologous sequences is
selected. The study demonstrates that substitution of a
surface exposed amino acid has the capacity to modify the
binding characteristics of a monoclonal antibody, which
is not surprising considering common knowledge. The

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experiments described are not designed to assess modulation in the binding of polyclonal antibodies such as allergic patients' serum IgE. One of the experiments contained do apply serum IgE and although this experiment is not suitable for quantitative assessment, IgE binding does not seem to be affected by the mutations performed.

The article by Smith et al. (ref. 5) describes the use of site directed mutagenesis for the purpose of monoclonal antibody epitope mapping and reduction of IgE binding. The authors have no knowledge of the tertiary structure and make no attempt to assess the conservation of α -carbon backbone tertiary structure. The algorithm used does not ensure that amino acids selected for mutation are actually exposed to the molecular surface. Only one of the mutants described lead to a substantial reduction in IgE binding. This mutant is deficient in binding of all antibodies tested indicating that the tertiary structure is disrupted. The authors do not define a therapeutic strategy and initiation of a new immune response is not mentioned.

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The article by Colombo et al. (ref. 14) describes the study of an IgE binding epitope by use of site directed mutagenesis and peptide synthesis. The authors use a three dimensional computer model structure based on the crystal structure of a homologous protein to illustrate the presence of the epitope on the molecular surface. The further presence of an epitope on a different allergen showing primary structure homology is addressed using synthetic peptides representing the epitope. The therapeutic strategy is based on treatment using this synthetic peptide representing a monovalent IgE binding epitope. Conserved surface areas between homologous allergens as well as the therapeutic concept of initiating a new protective immune response are not

mentioned.

The article by Spangfort et al. (ref. 15) describes the three-dimensional structure and conserved surface exposed 5 patches of the major birch allergen. The article does not mention major IgE binding epitopes nor site directed mutagenesis, neither is therapeutic application addressed.

- In none of the studies described above is IgE binding reduced by substitution of surface exposed amino acids while conserving α -carbon backbone tertiary structure. The rationale behind above-mentioned approaches does not include the concept of dominant IgE binding epitopes and
- it does not include the therapeutic concept of initiating a new protective immune response.

BRIEF DESCRIPTION OF THE FIGURES

Figure 1 shows mutant-specific oligonucleotide primers used for Bet v 1 mutant number 1. Mutated nucleotides are underlined.

Figure 2 shows two generally applicable primers (denoted 25 "all-sense" and "all non-sense"), which were synthesised and used for all mutants.

Figure 3 shows an overview of all $Bet \ v \ 1$ mutations.

- 60 Figure 4 shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Glu45Ser mutant.
- 35 Figure 5 shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IqE from a pool

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of allergic patients by non-biotinylated Bet v 1 and by Ret v 1 mutant Asn28Thr+Lvs32Gln.

Figure 6 shows the inhibition of the binding of 5 biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Pro108Gly mutant.

Figure 7 shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Glu60Ser mutant.

Figure 8 shows the CD spectra of recombinant and 5 Triple-patch mutant, recorded at close to equal concentrations.

Figure 9 shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Triple-patch mutant.

Figure 10 shows solvent accessibility of individually aligned antigen 5 residues and alignment of Vespula antigen 5 sequences (left panel). On the right panel of Figure 10 is shown the molecular surface of antigen 5 with conserved areas among Vespula antigen 5:s.

Figure 11 shows the sequence of the primer corresponding to the amino terminus of Ves v 5 derived from the sense strand. The sequence of the downstream primer is derived from the non-sense strand.

Figure 12 shows two generally applicable primers (denoted "all sense" and "all non-sense", which were synthesised and used for all mutants.

Figure 13 shows an overview of all Ves v 5 mutations.

Figure 14 shows the inhibition of the binding of biotinylated recombinant Ves v 5 to serum IgE from a pool of allergic patients by non-biotinylated Ves v 5 and by Ves v 5 Lys72Ala mutant.

OBJECT OF THE INVENTION

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Rationale behind the present invention

The current invention is based on a unique rationale. According to this rationale the mechanism of successful 15 allergy vaccination is not an alteration of the ongoing Th2-type immune response, but rather a parallel initiation of a new Th1-type immune response involving tertiary epitope recognition by B-cells and antibody formation. This model is supported by the observation 20 that levels of specific IgE are unaffected by successful vaccination treatment, and that successful treatment is often accompanied by a substantial rise in allergen specific IgG4. In addition, studies of nasal biopsies before and after allergen challenge do not show a 25 reduction in T cells with the Th2-like phenotype, but rather an increase in Th1-like T cells are observed. When vaccine (or pharmaceutical compositions) administered through another route than the airways, it is hypothesised, that the new Th1-like immune response 30 evolves in a location physically separated from the ongoing Th2 response thereby enabling the two responses to exist in parallel.

Another important aspect of the rationale behind the 35 current invention is the assertion of the existence of dominant IgE binding epitopes. It is proposed that these 10

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dominant IgE binding epitopes are constituted by tertiary structure dependent coherent surface areas large enough to accommodate antibody binding and conserved among isoallergens, variants, and/or homologous allergens from related species. The existence of cross-reactive IgE capable of binding similar epitopes on homologous allergens is supported by the clinical observation that allergic patients often react to several closely related species, e.g. alder, birch, and hazel, multiple grass species, or several species of the house dust mite genus Dermatophagoides. It is furthermore supported laboratory experiments demonstrating IgE cross-reactivity between homologous allergens from related species and the capacity of one allergen to inhibit the binding of IgE to homologous allergens (Ipsen et al. 1992, ref. 16). It is 15 well known that exposure and immune responses are related in a dose dependent fashion. Based on the combination of these observations it is hypothesised that conserved surface areas are exposed to the immune system in higher doses than non-conserved surface areas resulting in the generation of IgE antibodies with higher affinities, hence the term 'dominant IgE binding epitopes'.

According to this rationale it is essential that the allergen has an α -carbon backbone tertiary structure which essentially is the same as that of the natural allergen, thus ensuring conservation of the surface areas surrounding conserved patches topology of representing targets for mutagenesis aimed at reducing IgE binding. By fulfilling these criteria the allergen has the potential to be administered in relatively higher doses improving its efficacy in generating a protective immune response without compromising safety.

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The present invention relates to the introduction of artificial amino acid substitutions into defined critical positions while retaining the α -carbon backbone tertiary structure of the allergen.

The invention provides a recombinant allergen, which is a non-naturally occurring mutant derived from a naturally occurring allergen, wherein at least one surface-exposed, conserved amino acid residue of a B cell epitope is substituted by another residue which does not occur in the same position in the amino acid sequence of any known homologous protein within the taxonomic order from which said naturally occurring allergen originates, said mutant allergen having essentially the same $\alpha\text{-carbon}$ backbone tertiary structure as said naturally occurring allergen, and the specific IgE binding to the mutated allergen being reduced as compared to the binding to said naturally occurring allergen.

20 Such recombinant allergen is obtainable by

- a) identifying amino acid residues in a naturally occurring allergen which are conserved with more than 70% identity in all known homologous proteins within the taxonomic order from which said naturally occurring allergen originates;
- b) defining at least one patch of conserved amino acid residues being coherently connected over at least 400 Å of the surface of the three-dimensional of the allergen molecule as defined by having a solvent accessibility of at least 20%, said at least one patch comprising at least one B cell epitope; and
- 35 c) substituting at least one amino acid residue in said at least one patch by another amino acid being non-con-

servative in the particular position while essentially preserving the overall α -carbon backbone tertiary structure of the allergen molecule.

5 Specific IgE binding to the mutated allergen is preferably reduced by at least 5%, preferably at least 10% in comparison to naturally-occurring isoallergens or similar recombinant proteins in an immuno assay with sera from scource-specific IgE reactive allergic patients or 10 pools thereof.

Recombinant allergens according to the invention may suitably be derived from inhalation allergens originating i.a. from trees, grasses, herbs, fungi, house dust mites, 15 cockroaches and animal hair and dandruff. Important pollen allergens from trees, grasses and herbs are such originating from the taxonomic orders of Fagales, Oleales and Pinales including i.a. birch (Betula), alder (Alnus), hazel (Corylus), hornbeam (Carpinus) and olive (Olea), 20 the order of Poales including i.a. grasses of the genera Lolium, Phelum, Poa, Cynodon, Dactylis and Secale, the orders of Asterales and Urticales including i.a. herbs of the genera Ambrosia and Artemisia. Important inhalation allergens from fungi are i.a. such originating from the 25 genera Alternaria and Cladosporium. Other important inhalation allergens are those from house dust mites of the genus Dermatophagoides, those from cockroaches and those from mammals such as cat, dog and horse. Further, recombinant allergens according to the invention may be derived from venom allergens including such originating from stinging or biting insects such as those from the Hymenoptera including taxonomic order of (superfamily Apidae), wasps (superfamily Vespidea), and ants (superfamily Formicoidae).

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Specific allergen components include e.g. Bet v 1 (B.

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verrucosa, birch), Aln g 1 (Alnus glutinosa, alder), Cor a 1 (Corylus avelana, hazel) and Car b 1 (Carpinus betulus, hornbeam) of the Fagales order. Others are Cry j 1 (Pinales), Amb a 1 and 2, , Art v 1 (Asterales), Par j 5 1 (Urticales), Ole e 1 (Oleales), Ave e 1, Cyn d 1, Dac q 1, Fes p 1, Hol 1 1, Lol p 1 and 5, Pas n 1, Phl p 1 and 5, Poa p 1, 2 and 5, Sec c 1 and 5, and Sor h 1 (various grass pollens), Alt a 1 and Cla h 1 (fungi), Der f 1 and 2, Der p 1 and 2 (house dust mites, D. farinae and D. 10 pteronyssinus, respectively), Bla g 1 and 2, Per a 1 germanica and Periplaneta (cockroaches, Blatella americana, respectively), Fel d 1 (cat), Can f 1 (dog), Equ c 1, 2 and 3 (horse), Apis m 1 and 2 (honeybee), Ves q 1, 2 and 5, Pol a 1, 2 and 5 (all wasps) and Sol i 1, 15 2, 3 and 4 (fire ant).

In one embodiment, the recombinant allergen is derived from Bet v 1. Examples of substitutions are Thr10Pro, Asp25Gly, (Asn28Thr + Lys32Gln), Glu45Ser, Asn47Ser, 290 Lys55Asn, Thr77Ala, Pro108Gly and (Asn28Thr, Lys32Gln, Glu45Ser, Pro108Gly. As apparent, the recombinant allergens may have one or more substitutions.

In another embodiment, the recombinant allergen is 25 derived from a venom allergen from the taxonomic order of Vespidae, Apidae and Formicoidae.

In a further embodiment, the recombinant allergen is derived from Ves v 5. Examples of substitutions are Uss72Ala and Tyr96Ala. As apparent, the recombinant allergens may have one or more substitutions.

The present invention also provides a method of preparing a recombinant allergen as defined herein, comprising

a) identifying amino acid residues in a naturally

occurring allergen which are conserved with more than 70% identity in all known homologous proteins within the taxonomic order from which said naturally occurring allergen originates;

b) defining at least one patch of conserved amino acid residues being coherently connected over at least 400 Å² of the surface of the three-dimensional structure of the allergen molecule as defined by having a solvent accessibility of at least 20 %, said at least one patch comprising at least one B cell epitope, and

c) substituting at least one amino acid residue in said at least one patch by another amino acid being non-conservative in the particular position while essentially preserving the overall α -carbon backbone tertiary structure of the allergen molecule.

In this method the best results are obtained by ranking the amino acid residues of said at least one patch with respect to solvent accessibility and substituting one or more amino acids among the more solvent accessible ones.

Generally, in the method according to the invention the substitution of one or more amino acid residues in said B cell epitope or said at least one patch is carried out by site-directed mutagenesis.

Conservation of α -carbon backbone tertiary structure is best determined by obtaining identical structures by x-ray crystallography or NMR before and after mutagenesis. In absence of structural data describing the mutant indistinguishable CD-spectra or immunochemical data, e.g. antibody reactivity, may render conservation of α -carbon backbone tertiary structure probable, if compared to the data obtained by analysis of a structurally determined

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molecule.

Further, the present invention provides a pharmaceutical composition comprising a recombinant allergen as defined herein in combination with a pharmaceutically acceptable carrier and/or excipient, and optionally an adjuvant.

Such pharmaceutical composition may be in the form of a vaccine against allergic reactions elicited by a naturally occurring allergen in patients suffering from allergy.

In a further aspect, the present invention relates to a method of generating an immune response in a subject, which method comprises administering to the subject at least one recombinant allergen as defined herein, or a pharmaceutical composition comprising at least one recombinant allergen as defined herein.

The pharmaceutical composition of the invention can be prepared by a process comprising mixing at least one recombinant allergen as defined herein with pharmaceutically acceptable substances and/or excipients.

In a particular embodiment, the present invention concerns the vaccination or treatment of a subject, which vaccination of treatment comprises administering to the subject at least one recombinant allergen as defined herein or a pharmaceutical composition as defined herein.

The pharmaceutical compositions of the invention are obtainable by the process defined above.

In another embodiment, the recombinant allergens of the invention are suitable for use in a method for the treatment, prevention or alleviation of allergic reactions, such method comprising administering to a subject a recombinant allergen as defined herein or a pharmaceutical composition as defined herein.

DETAILED DESCRIPTION OF THE INVENTION

Criteria for substitution

For molecules for which the tertiary structure has been to determined (e.g. by x-ray crystallography, or NMR electron microscopy), the mutant carrying the substituted amino acid(s) should preferably fulfil the following criteria:

- 15 1. The overall α -carbon backbone tertiary structure of the molecule should be conserved. Conserved is defined as an average root mean square deviation of the atomic coordinates comparing the structures below 2A. This is important for two reasons: a) It is anticipated that the entire surface of the natural allergen constitutes an overlapping continuum of potential antibody-binding epitopes. The majority of the surface of the molecule is not affected by the substitution(s), and thus retain its antibody-binding properties, which is important for the generation of new protective antibody specificities being directed at epitopes present also on the natural allergen. b) Stability, both concerning shelf-life and upon injection into body fluids.
- 2. The amino acid(s) to be substituted should be located at the surface, and thus be accessible for antibody-binding. Amino acids located on the surface are defined as amino acids in the three-dimensional structure having a solvent (water) accessibility of at least 20%, suitably 30-80%, more suitably 30-80%. Solvent accessibility is defined as the area of the molecule accessible to a

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sphere with a radius comparable to a solvent (water, r = 1.4 Å) molecule.

- 3. The substituted amino acid(s) should be located in conserved patches larger than 400 Å². Conserved patches are defined as coherently connected areas of surface exposed conserved amino acid residues and backbone. Conserved amino acid residues are defined by sequence alignment of all known (deduced) amino acid sequences of homologues proteins within the taxonomical order. Amino acid positions having identical amino acid residues in more than 90% of the sequences are considered conserved. Conserved patches are expected to contain epitopes to which the IgE of the majority of patients is directed.
 - 4. Within the conserved patches amino acids for mutagenesis should preferentially be selected among the most solvent (water) accessible ones located preferably near the centre of the conserved patch.

Preferentially, a polar amino acid residue is substituted by another polar residue, and a non-polar amino acid residue is substituted by another non-polar residue.

Preparation of vaccines is generally well-known in the 25 art. Vaccines are typically prepared as injectables either as liquid solutions or suspensions. Such vaccine may also be emulsified or formulated so as to enable nasal administration. The immunogenic component question (the recombinant allergen as defined herein) may 30 mixed with excipients which pharmaceutically acceptable and further compatible with the active ingredient. Examples of suitable excipients are water, saline, dextrose, glycerol, ethanol and the like as well as combinations thereof. The vaccine may additionally contain other substances such as wetting agents, emulsifying agents, buffering agents or adjuvants enhancing the effectiveness of the vaccine.

Vaccines are most frequently administered parenterally by 5 subcutaneous or intramuscular injection. Formulations which are suitable for administration by another route include oral formulations and suppositories. Vaccines for oral administration may suitably be formulated with excipients normally employed for such formulations, e.g. pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharine, cellulose, magnesium carbonate and the like. The composition can be formulated as solutions, suspensions, emulsions, tablets, sustained formulations, capsules, release aerosols, powders, or granulates.

The vaccines are administered in a way so as to be compatible with the dosage formulation and in such amount as will be therapeutically effective and immunogenic. The quantity of active component contained within the vaccine depends on the subject to be treated, i.a. the capability of the subject's immune system to respond to the treatment, the route of administration and the age and weight of the subject. Suitable dosage ranges can vary within the range from about 0.0001 µg to 1000 µg.

As mentioned above, an increased effect may be obtained by adding adjuvants to the formulation. Examples of such adjuvants are aluminum hydroxide and phosphate (alum) as 30 a 0.05 to 0.1 percent solution in phosphate buffered saline, synthetic polymers of sugars used as 0.25 percent solution. Mixture with bacterial cells such as C. parvum, endotoxins or lipopolysaccharide components of gramnegative bacteria, emulsion in physiologically acceptable oil vehicles such as mannide monoaleate (Aracel A) or emulsion with 20 percent solution of a perfluorocarbon

(e.g. Fluosol-DA) used as a block substitute may also be employed. Other adjuvants such as Freund's complete and incomplete adjuvants as well as QuilA and RIBI may also be used.

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Most often, multiple administrations of the vaccine will be necessary to ensure an effect. Frequently, the vaccine is administered as an initial administration followed by subsequent inoculations or other administrations. The 10 number of vaccinations will typically be in the range of from 1 to 50, usually not exceeding 35 vaccinations. Vaccination will normally be performed from biweekly to bimonthly for a period of 3 months to 5 years. This is contemplated to give desired level of prophylactic or 15 therapeutic effect.

The recombinant allergen may be used as a pharmaceutical preparation, which is suitable for providing a certain protection against allergic responses during the period of the year where symptoms occur (prophylaxis). Usually, the treatment will have to be repeated every year to maintain the protective effect. Preparations formulated for nasal application are particular suited for this purpose.

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The present invention is further illustrated by the following non-limiting examples.

EXAMPLES

EXAMPLE 1

Identification of common epitopes within Fagales pollen allergens

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The major birch pollen allergen Bet v 1 shows about 90%

amino acid sequence identity with major allergens from pollens of taxonomically related trees, i.e Fagales (for instance hazel and hornbeam) and birch pollen allergic patients often show clinical symptoms of allergic cross-reactivity towards these Bet v 1 homologous proteins.

Bet v 1 also shows about 50-60% sequence identity with allergic proteins present in certain fruits (for instance apple and cherry) and vegetables (for instance celery and carrot) and there are clinical evidence for allergic cross-reactivity between Bet v 1 and these food related proteins.

In addition, Bet v 1 shares significant sequence identity
15 (20-40%) with a group of plant proteins called
pathogenesis-related proteins (PR-10), however there are
no reports of allergic cross-reactivity towards these PR10 proteins.

Molecular modelling suggests that the structures of Fagales and food allergens and PR-10 proteins are close to be identical with the Bet v l structure.

The structural basis for allergic Bet v 1 crossreactivity was reported in (Gajhede et al 1996, ref. 17)
where three patches on the molecular surface of Bet v 1
could be identified to be common for the known major tree
pollen allergens. Thus, any IgE recognising these patches
on Bet v 1 would be able to cross-react and bind to other
fagales major pollen allergens and give rise to allergic
symptoms. The identification of these common patches was
performed after alignment of all known amino acid
sequences of the major tree pollen allergens in
combination with an analysis of the molecular surface of
Bet v 1 revealed by the α-carbon backbone tertiary
structure reported in ref. 17. In addition, the patches

were defined to have a certain minimum size (>400 ${\rm \AA}^2$) based on the area covered by an antibody upon binding.

Selection of amino acid residues for site-directed mutagenesis

Amino acid residues for site-directed mutagenesis were selected among residues present in Bet v 1 specific areas and the common patches since modifications of these is expected to affect the binding of serum IgE from the majority of patients showing clinical tree pollen allergic cross-reactivity.

The relative orientation and percentage of solvent-exposure of each amino acid residue within respective patch was calculated based on their atomic coordinates. Residues having a low degree of solvent exposure (<20%) were not regarded relevant for mutagenesis due to the possible disruption of the structure or lack of antibody interaction. The remaining residues were ranked according to their degree of solvent-exposure.

Sequence alignment

25 Sequences homologous to the query sequence (Bet v 1 No. 2801, WHO IUIS Nomenclature Subcommittee on Allergens) were derived from GenBank and EMBL sequence databases by a BLAST search (Altschul et al., ref. 18). All sequences with BLAST reported probabilities less than 0.1 were taken into consideration and one list were constructed containing a non-redundant list of homologous sequences. These were aligned by CLUSTAL W (Higgins et al., ref. 19) and the percentage identity were calculated for each position in the sequence considering the complete list or taxonomically related species only. A total of 122 sequences were homologous to Bet v 1 No. 2801 of which 57

sequences originates from taxonomically related species.

Cloning of the gene encoding Bet v 1

5 RNA was prepared from Betula verrucosa pollen (Allergon, Sweden) by phenol extraction and LiCl precipitation. Oligo (dT)-cellulose affinity chromatography was performed batch-wise in Eppendorph tubes, and double-stranded cDNA was synthesised using a commercially available kit (Amersham). DNA encoding Bet v l was amplified by PCR and cloned. In brief, PCR was performed using cDNA as template, and primers designed to match the sequence of the cDNA in positions corresponding to the amino terminus of Bet v l and the 3'-untranslated region, respectively.

15 The primers were extended in the 5'-ends to accommodate restriction sites (NcoI and HindIII) for directional cloning into pKK233-2.

Subcloning into pMAL-c

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The gene encoding Bet v 1 was subsequently subcloned into the maltose binding protein fusion vector pMAL-c (New England Biolabs). The gene was amplified by PCR and subcloned in frame with malE to generate maltose binding protein (MBP)-Bet v 1 protein fusion operons in which MBP and $Bet \ v \ 1$ were separated by a factor X_a protease clevage site positioned to restore the authentic aminoterminal sequence of Bet v 1 upon cleavage, as described in ref. 15. In brief, PCR was performed using 30 pKK233-3 with Bet v 1 inserted as template and primers corresponding to the amino- and carboxyterminus of the protein, respectively. The promoter proximal primer was extended in the 5'-end to accommodate 4 codons encoding an in frame factor Xa protease cleavage site. Both primers were furthermore extended in the 5'-ends to accommodate restriction sites (KpnI) for cloning.

Bet v 1 encoding genes were subcloned using 20 cycles of PCR to reduce the frequency of PCR artefacts.

In vitro mutagenesis

In vitro mutagenesis was performed by PCR using recombinant pkAL-c with $Bet\ v\ 1$ inserted as template. Each mutant $Bet\ v\ 1$ gene was generated by 3 PCR reactions using 4 primers.

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Two mutation-specific oligonucleotide primers were synthesised accommodating each mutation, one for each DNA strand, see Figs. 1 and 2, Using the mutated nucleotide(s) as starting point both primers were extended 7 nucleotides in the 5'-end and 15 nucleotides in the 3'-end. The extending nucleotides were identical in sequence to the $Bet\ v\ I$ gene in the actual region.

Two generally applicable primers (denoted "all-sense" and "all non-sense" in Figure 2) were furthermore synthesised and used for all mutants. These primers were 15 nucleotides in length and correspond in sequence to regions of the pMAL-c vector approximately 1 kilobase upstream and downstream from the Bet v 1. The sequence of the upstream primer is derived from the sense strand and the sequence of the downstream primer is derived from the non-sense strand, see Fig. 2.

Two independent PCR reactions were performed essentially according to standard procedures (Saiki et al 1988, ref. 20) with the exception that only 20 temperature cycles were performed in order to reduce the frequency of PCR artefacts. Each PCR reaction used pMAL-c with Bet v l inserted as template and one mutation-specific and one generally applicable primer in meaningful combinations.

Introduction of the four amino acid substitutions (Asn28Thr, Lys32Gln, Glu45Ser, Pro108Gly) in the Triple-patch mutant were performed like described above in a step by step process. First the Glu45Ser mutation then the Pro108Gly mutation and last the Asn28Thr, Lys32Gln mutations were introduced using pMAL-c with inserted Bet v 1 No. 2801, Bet v 1 (Glu45Ser), Bet v 1 (Glu45Ser, Pro108Gly) as templates, respectively.

The PCR products were purified by agarose gel electrophoresis and electro-elution followed by ethanol precipitation. A third PCR reaction was performed using the combined PCR products from the first two PCR reactions as template and both generally applicable primers. Again, 20 cycles of standard PCR were used. The PCR product was purified by agarose gel electrophoresis and electro-elution followed by ethanol precipitation, cut with restriction enzymes (BsiWI/EcoRI), and ligated directionally into pMAL-c with Bet v I inserted presented to restricted with the same enzymes.

Figure 3 shows an overview of all 9 Bet v 1 mutations, which are as follows

25 Thr10Pro, Asp25Gly, Asn28Thr + Lys32Gln, Glu45Ser, Asn47Ser, Lys55Asn, Glu60Ser (non-patch), Thr77Ala and Pro108Gly. An additional four mutant with four mutations was also prepared (Asn28Thr, Lys32Gln, Glu45Ser, Pro108Gly). Of these, five were selected for further testing: Asn28Thr + Lys32Gln, Glu45Ser, Glu60Ser, Pro108Gly and the Triple-patch mutant Asn28Thr, Lys32Gln, Glu45Ser, Pro108Gly.

Nucleotide sequencing

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Determination of the nucleotide sequence of the Bet v 1

encoding gene was performed before and after subcloning, and following in vitro mutagenesis, respectively.

Plasmid DNA's from 10 ml of bacterial culture grown to saturation overnight in LB medium supplemented with 0.1 g/l ampicillin were purified on Qiagen-tip 20 columns and sequenced using the Sequenase version 2.0 DNA sequencing kit (USB) following the recommendations of the suppliers.

Expression and purification of recombinant Bet v 1 and mutants

Recombinant Bet v 1 (Bet v 1 No. 2801 and mutants) were over-expressed in Escherichia coli DH 5a fused to maltose-binding protein and purified as described in ref. 15. Briefly, recombinant E.coli cells were grown at 37°C to an optical density of 1.0 at 436 nm, whereupon expression of the Bet v 1 fusion protein was induced by addition of IPTG. Cells were harvested by centrifugation 20 3 hours post-induction, re-suspended in lysis buffer and broken by sonication. After sonication and additional centrifugation, recombinant fusion protein was isolated by amylose affinity chromatography and subsequently cleaved by incubation with Factor Xa (ref. 15). After F Xa cleavage, recombinant Bet v 1 was isolated by 25 gelfiltration and if found necessary, subjected to another round of amylose affinity chromatography in order to remove trace amounts of maltose-binding protein.

30 Purified recombinant Bet v 1 was concentrated by ultrafiltration to about 5 mg/ml and stored at 4 °C. The final yields of the purified recombinant Bet v 1 preparations were between 2-5 mg per litre E. coli cell culture.

The purified recombinant Bet v 1 preparations appeared as

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single bands after silver-stained SDS-polyacrylamide electrophoresis with an apparent molecular weight of 17.5 kDa. N-terminal sequencing showed the expected sequences as derived from the cDNA nucleotide sequences and 5 quantitative amino acid analysis showed the expected amino acid compositions.

We have previously shown (ref. 15) that recombinant Bet v 1 No. 2801 is immunochemically indistinguishable from naturally occurring Bet v 1.

Immunoelectrophoresis using rabbit polyclonal antibodies

The seven mutant Bet v 1 were produced as recombinant Bet 15 v 1 proteins and purified as described above and tested for their reactivity towards polyclonal rabbit antibodies raised against Bet v 1 isolated from birch pollen. When analysed by immunoelectrophoresis (rocket-line immunoelectrophoresis) under native conditions, the 20 rabbit antibodies were able to precipitate all mutants, indicating that the mutants had conserved α-carbon backbone tertiary structure.

These results suggested that non-naturally occurring substitutions introduced on the molecular surface of Bet v 1 can reduce a polyclonal antibody response raised against naturally occurring Bet v 1 without distortion of the overall α-carbon backbone tertiary allergen structure. In order to analyse the effect on human polyclonal IgE-response, the mutants Glu45Ser, Pro108Gly, Asn28Thr+Lys32Gln and Glu60Ser were selected for further analysis.

Bet v 1 Glu45Ser mutant

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Glutamic acid in position 45 show a high degree of

solvent-exposure (40%) and is located in a molecular surface patch common for Fagales allergens (patch I). A serine residue was found to occupy position 45 in some of the Bet v 1 homologous PR-10 proteins arguing for that glutamic acid can be replaced by serine without distortion of the α-carbon backbone tertiary structure. In addition, as none of the known Fagales allergen sequences have serine in position 45, the substitution of glutamic acid with serine gives rise to a non-naturally occurring Bet v 1 molecule.

T cell proliferation assay using recombinant Glu45Ser Bet v 1 mutant

15 The analysis was carried out as described in Spangfort et al 1996a. It was found that recombinant Bet v 1 Glu45Ser mutant was able to induce proliferation in T cell lines from three different birch pollen allergic patients with stimulation indices similar to recombinant and naturally 20 occurring.

<u>Crystallisation</u> and structural determination of recombinant Glu45Ser Bet v 1

crystals of recombinant Glu45Ser Bet v 1 were grown by vapour diffusion at 25°C, essentially as described in (Spangfort et al 1996b, ref. 21). Glu45Ser Bet v 1, at a concentration of 5 mg/ml, was mixed with an equal volume of 2.0 M ammonium sulphate, 0.1 M sodium citrate, 1% (v/v) dioxane, pH 6.0 and equilibrated against 100x volume of 2.0 M ammonium sulfate, 0.1 M sodium citrate, 1% (v/v) dioxane, pH 6.0. After 24 hours of equilibration, crystal growth was induced by applying the seeding technique described in ref. 21, using crystals of recombinant wild-type Bet v 1 as a source of seeds.

After about 2 months, crystals were harvested and analysed using X-rays generated from a Rigaku rotating anode as described in ref. 21 and the structure was solved using molecular replacement.

Structure of Bet v 1 Glu45Ser mutant

The structural effect of the mutation was addressed by growing three-dimensional Bet v 1 Glu45Ser protein crystals diffracting to 3.0 Å resolution when analysed by X-rays generated from a rotating anode. The substitution of glutamic acid to serine in position 45 was verified by the Bet v 1 Glu45Ser structure electron density map which also showed that the overall α-carbon backbone tertiary structure is preserved.

IqE-binding properties of Bet v 1 Glu45Ser mutant

The IgE-binding properties of Bet v 1 Glu45Ser mutant was compared with recombinant Bet v 1 in a fluid-phase IgE-inhibition assay using a pool of serum IgE derived from birch alleraic patients.

Recombinant Bet v 1 no. 2801 was biotinylated at a molar ratio of 1:5 (Bet v 1 no. 2801:biotin). The inhibition assay was performed as follows: a serum sample (25 µl) was incubated with solid phase anti IgE, washed, resuspended and further incubated with a mixture of biotinylated Bet v 1 no. 2801 (3.4 nM) and a given mutant (0-28.6 nM). The amount of biotinylated Bet v 1 no. 2801 bound to the solid phase was estimated from the measured RLU after incubation with acridinium ester labelled streptavidin. The degree of inhibition was calculated as the ratio between the RLU's obtained using buffer and mutant as inhibitor.

Figure 4 shows the inhibition of the binding of biotinylated recombinant $Bet \ v \ 1$ to serum IgE from a pool of allergic patients by non-biotinylated $Bet \ v \ 1$ and by $Ref \ v \ 1$ Glu4SSer mutant.

There is a clear difference in the amount of respective recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant Bet v 1 reaches 50% inhibition at about 6.5 on gwhereas the corresponding concentration for Bet v 1 Glu45Ser mutant is about 12 ng. This show that the point mutation introduced in Bet v 1 Glu45Ser mutant lowers the affinity for specific serum IgE by a factor of about 2. The maximum level of inhibition reached by the Bet v 1 Glu45Ser mutant is clearly lower compared to recombinant Bet v 1. This may indicate that after the Glu45Ser substitution, some of the specific IgE present in the serum pool are unable to recognise the Bet v 1 Glu45Ser mutant.

Bet v 1 mutant Asn28Thr+Lys32Gln

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Aspartate and lysine in positions 28 and 32, respectively show a high degree of solvent-exposure (35% and 50%, 25 respectively) and are located in a molecular surface patch common for Fagales allergens (patch II). In the structure, aspartate 28 and lysine 32 are located close to each other on the molecular surface and most likely interact via hydrogen bonds. A threonine and a gluatamate residue were found to occupy positions 28 and 32, respectively in some of the Bet v 1 homologous PR-10 proteins arguing for that aspartate and lysine can be replaced with threonine and glutamate, respectively without distortion of the α-carbon backbone tertiary structure. In addition, as none of the naturally occurring isoallergen sequences have threonine and

glutamate in positions 28 and 32, respectively, the substitutions gives rise to a non-naturally occurring Bet v 1 molecule.

5 IgE-binding properties of Bet v 1 mutant Asn28Thr+Lys32Gln

The IgE-binding properties of mutant Asn28Thr+Lys32Gln was compared with recombinant Bet v 1 in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.

Figure 5 shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool is of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 mutant Asn28Thr+Lys32Gln.

There is a clear difference in the amount of respective recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant Bet v 1 reaches 50% inhibition at about 6.5 ng whereas the corresponding concentration for Bet v 1 mutant Asn28Thr+Lys32Gln is about 12 ng. This show that the point mutations introduced in Bet v 1 mutant Asn28Thr+Lys32Gln lowers the affinity for specific serum IgE by a factor of about 2.

The maximum level of inhibition reached by the Bet v 1 mutant Asn28Thr+Lys32Gln mutant is clearly lower compared to recombinant Bet v 1. This may indicate that after the Asn28Thr+Lys32Gln substitutions, some of the specific IgE present in the serum pool are unable to recognise the Bet v 1 mutant Asn28Thr+Lys32Gln.

35 Bet v 1 mutant Pro108Gly

Proline in position 108 show a high degree of solvent-exposure (60%) and is located in a molecular surface patch common for Fagales allergens (patch III). A glycine residue was found to occupy position 108 in some of the 5 Bet ν 1 homologous PR-10 proteins arguing for that proline can be replaced with glycine without distortion of the α -carbon backbone tertiary structure. In addition, as none of the naturally occurring isoallergen sequences have glycine in position 108, the substitution of proline 0 with glycine gives rise to a non-naturally occurring Bet ν 1 molecule.

IgE-binding properties of Bet v 1 Pro108Gly mutant

- The IgE-binding properties of Bet v 1 Pro108Gly mutant was compared with recombinant Bet v 1 in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.
- Figure 6 shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Pro108Gly mutant.
- There is a clear difference in the amount of respective recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant Bet v 1 reaches 50% inhibition at about 6.5 ng whereas the corresponding concentration for Bet v 1 Prol08Gly is 15 ng. This show that the single point mutation introduced in Bet v 1 Prol08Gly lowers the affinity for specific serum IgE by a factor of about 2.
- The maximum level of inhibition reached by the Bet v 1
 Fro108Gly mutant is somewhat lower compared to recombinant Bet v 1. This may indicate that after the

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Prol08Gly substitution, some of the specific IgE present in the serum pool are unable to recognise the Bet v 1 Prol08Gly mutant.

5 Bet v 1 mutant Glu60Ser (non-patch mutant)

Glutamic acid in position 60 show a high degree of solvent-exposure (60%) however, it is not located in a molecular surface patch common for Fagales allergens. A 10 serine residue was found to occupy position 60 in some of the Bet v l homologous PR-10 proteins arguing for that glutamic acid can be replaced with serine without distortion of the α-carbon backbone tertiary structure. In addition, as none of the naturally occurring isoallergen sequences have serine in position 60, the substitution of glutamic acid with serine gives rise to a non-naturally occurring Bet v l molecule.

IgE-binding properties of Bet v 1 Glu60Ser mutant

The IgE-binding properties of $Bet\ v\ 1$ Glu60Ser mutant was compared with recombinant $Bet\ v\ 1$ in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.

Figure 7 shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Glu60Ser mutant. In contrast to the Glu45Ser, Pro108Gly and Asn28Thr+Lys32Gln mutants, the substitution glutamic acid 60 to serine, does not shown any significant effect on the IgE-binding properties of. This indicates that substitutions outside the defined Fagales common patches only have a marginal effect on the binding of specific serum IgE supporting the concept that conserved allergen molecular surface areas harbours

dominant IgE-binding epitopes.

Bet v 1 Triple-patch mutant

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5 In the Triple-patch mutant, the point mutations (Glu45Ser, Asn28Thr+1ys32Gln and Pro108Gly) introduced in the three different common Fagales patches, described above, were simultaneously introduced in creating an artificial mutant carrying four amino acid substitutions.

Structural analysis of Bet v 1 Triple-patch mutant

The structural integrity of the purified Triple-patch mutant was analysed by circular dichroism (CD) 15 spectroscopy. Figure 8 shows the CD spectra of recombinant and Triple-patch mutant, recorded at close to equal concentrations. The overlap in peak amplitudes and positions in the CD spectra from the two recombinant proteins shows that the two preparations contain equal amounts of secondary structures strongly suggesting that the α-carbon backbone tertiary structure is not affected by the introduced amino acid substitutions.

IgE-binding properties of Bet v 1 Triple-patch mutant

The IgE-binding properties of Bet v 1 Triple-patch mutant was compared with recombinant Bet v 1 in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.

Figure 9 shows the inhibition of the binding of biotinylated recombinant Bet v 1 to serum IgE from a pool of allergic patients by non-biotinylated Bet v 1 and by Bet v 1 Triple-patch mutant. In contrast to the single mutants described above, the inhibition curve of the Triple-patch mutant is no longer parallel relative to

recombinant. This shows that the substitutions introduced in the Triple-patch mutant has changed the IgE-binding properties and epitope profile compared to recombinant. The lack of parallellity makes it difficult to quantify the decrease of the Triple-patch mutant affinity for specific serum IgE.

Recombinant Bet v 1 reaches 50% inhibition at about 6 ng whereas the corresponding concentration for Bet v 1 10 Triple-patch mutant is 30 ng, i.e a decrease in affinity by a factor 5. However, in order to reach 80% inhibition the corresponding values are 20 ng and 400 ng, respectively, i.e a decrease by a factor 20.

IS <u>T cell proliferation assay using recombinant Bet v 1</u> Triple-patch mutant

The analysis was carried out as described in ref. 15. It was found that recombinant Bet v 1 Triple-patch mutant 20 was able to induce proliferation in T cell lines from three different birch pollen allergic patients with stimulation indices similar to recombinant and naturally occurring. This suggests that the Triple-patch mutant can initiate the cellular immune response necessary for antibody production.

EXAMPLE 2

Identification of common epitopes within Vespula vulgaris venom major allergen antigen 5

Antigen 5 is one of the three vespid venom proteins, which are known allergens in man. The vespids include hornets, yellow-jacket and wasps. The other two known slergens of vespid venoms are phospholipase A: and hyaluronidase. Antigen 5 from Vespula vulgaris (Ves v 5)

has been cloned and expressed as recombinant protein in the yeast system (Monsalve et al. 1999, ref. 22). The three-dimensional crystal structure of recombinant Ves v 5 has recently been determined at 1.8 Å resolution (in preparation). The main features of the structure consist of four β -strands and four α -helices arranged in three stacked layers giving rise to a " α - β - α sandwich". The sequence identity between Antigen 5 homologous allergens from different Vespula species is about 90% suggesting presence of conserved molecular surface areas and B cell epitopes.

The presence and identification of common patches was performed after alignment of all known amino acid is sequences, as previously described for tree pollen allergens, of the Vespula antigen 5 allergens in combination with an analysis of the molecular surface of Antigen 5 revealed by the three-dimensional structure of Ves v 5. Figure 10 shows solvent accessibility of individually aligned antigen 5 residues and alignment of Vespula antigen 5 sequences (left panel). On the right panel of figure 10 is shown the molecular surface of antigen 5 with conserved areas among Vespula antigen 5:s coloured.

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Selection of amino acid residues for site-directed mutagenesis

Amino acid residues for site-directed mutagenesis were selected among residues present the patches common for Vespula since modifications of these is expected to affect the binding of serum IgE from the majority of patients showing clinical Vespula allergic cross-reactivity.

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The relative orientation and percentage of solvent-

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exposure of each amino acid residue within respective patch was calculated based on their atomic coordinates. Residues having a low degree of solvent exposure were not regarded suitable for mutagenesis due to the possible disruption of the structure or lack of antibody interaction. The remaining residues were ranked according to their degree of solvent-exposure.

Cloning of the gene encoding Ves v 5

Total RNA was isolated from venom acid glands of *Vespula* vulgaris vespids as described in (Fang et al. 1988, ref. 23).

15 First-strand cDNA synthesis, PCR amplification and cloning of the Ves v 5 gene was performed as described in (Lu et al. 1993, ref. 24)

Subcloning into pPICZaA

The gene encoding $Ves\ v\ 5$ was subsequently sub-cloned into the pPICZ αA vector (Invitrogen) for secreted expression of $Ves\ v\ 5$ in $Pichia\ pastoris$. The gene was amplified by PCR and sub-cloned in frame with the coding sequence for the α -factor secretion signal of $Saccharomyces\ cerevisiae$. In this construct the α -factor is cleaved off, in vivo, by the $Pichia\ pastoris\ Kex2$ protease system during secretion of the protein.

In brief PCR was performed using Ves v 5 as template and primers corresponding to the amino- and carboxyterminus of the protein, respectively. The primers were extended in the 5'-end to accommodate restriction sites for cloning, EcoRI and XbaI, respectively. Nucleotides encoding the Kex2 cleavage site was in this construct positioned 18 nucleotides upstream to the amino terminus

of the protein, resulting in the expression of Ves v 5 with six additional amino acids, Glu-Ala-Glu-Ala-Glu-Phe, at the amino terminus.

Insertion of pPICZαA-Ves v 5 into P. pastoris

The pPICZ α A vectors with the Ves v 5 gene inserted was linearised by Sac I restriction and inserted into the AOXI locus on the Pichia pastoris genome. Insertion was 10 performed by homologous recombination on Pichia pastoris KM71 cells following the recommendations of Invitrogen.

In vitro mutagenesis

- In vitro mutagenesis was performed by PCR using recombinant pFICZoA with Ves v 5 inserted as template. Each mutant Ves v 5 gene was generated by 3 PCR reactions using 4 primers.
- Two mutation-specific oligonucleotide primers were synthesised accommodating each mutation, one for each DNA strand, see Figures 11 and 12. Using the mutated nucleotide(s) as starting point both primers were extended 6-7 nucleotides in the 5'-end and 12-13 nucleotides in the 3'-end. The extending nucleotides were identical in sequence to the Ves v 5 gene in the actual region.
- Two generally applicable primers (denoted "all sense" and 30 "all non-sense" in Figure 12) were furthermore synthesised and used for all mutants. To insure expression of Ves v 5 mutants with authentic amino terminus, one primer corresponding to the amino terminus of the protein was extended in the 5´-end with a Xho I site. Upon insertion of the Ves v 5 mutant genes into the pPICZaA vector, the Kex2 protease cleavage site was

regenerated directly upstream to the amino terminus of Ves v 5. The second primer was corresponding in sequence to a region of the pPICZaA vector positioned approximately 300 bp downstream from the Ves v 5 gene. 5 The sequence of the primer corresponding to the amino terminus of Ves v 5 is derived from the sense strand and the sequence of the downstream primer is derived from the non-sense strand, see Figure 11.

Two independent PCR reactions were performed essentially according to standard procedures (Saiki et al 1988) with the exception that only 20 temperature cycles were performed in order to reduce the frequency of PCR artefacts. Each PCR reaction used pPICZaA with Ves v 5 inserted as template and one mutation-specific and one generally applicable primer in meaningful combinations.

The PCR products were purified by using "Concert, Rapid PCR Purification System" (Life Technologies). A third PCR reaction was performed using the combined PCR products from the first two PCR reactions as template and both generally applicable primers. Again, 20 cycles of standard PCR were used. The PCR product was purified with the "Concert, Rapid PCR Purification System" (Life Technologies), cut with restriction enzymes (XhoI/XbaI), and ligated directionally into pPICZQA vector restricted with the same enzymes. Figure 13 shows an overview of all Ves v 5 mutations.

30 Insertion of pPICZαA-Ves v 5 mutants into P. pastoris

The pPICZαA vectors with the Ves v 5 mutant genes inserted were linearised by Sac I restriction and inserted into the AOXI locus on the Pichia pastoris genome. Insertions were performed by homologous recombination on Pichia pastoris KM71 cells following the

recommendations of Invitrogen.

Nucleotide sequencing

5 Determination of the nucleotide sequence of the Ves v 5 encoding gene was performed before and after subcloning, and following in vitro mutagenesis, respectively.

Plasmid DNA's from 10 ml of bacterial culture grown to saturation overnight in LB medium supplemented with 0.1 g/l ampicillin were purified on Qiagen-tip 20 columns and sequenced using the Sequenase version 2.0 DNA sequencing kit (USB) following the recommendations of the suppliers.

15 Expression and purification of recombinant Ves v 5

Recombinant yeast cells of *Pichia pastoris* strain KM71 were grown in 500 ml bottles containing 100 ml of pH 6.0 phosphate buffer containing yeast nitrogen base, biotin, 20 glycerol and histidine at 30°C with orbital shaking at 225 rpm until A₅₀₀ nm of 4-6. Cells were collected by centrifugation and re-suspended in 10 ml of similar buffered medium containing methanol in place of glycerol. Incubation was continued at 30°C for 7 days with daily 25 addition of 0.05 ml methanol.

Cells were harvested by centrifugation and the collected culture fluid was concentrated by ultrafiltration. After dialysis against 50 mM ammonium acetate buffer, pH 4.6, the sample was applied to a FPLC (Pharmacia) SE-53 cation exchange column equilibrated in the same buffer. The column was eluated with a 0-1.0 M NaCl, 50 mM ammonium acetate linear gradient. The recombinant Ves v 5 peak eluting at about 0.4 M NaCl was collected and dialysed against 0.02 N acetic acid. After concentration to about 10 mg/ml, the purified Ves v 5 was stored at 4°C.

Crystallisation of recombinant Ves v 5

Crystals of Ves v 5 was grown by the vapour diffusion 5 technique at 25°C. For crystallisation, 5 µl of 5 mg/ml Ves v 5 was mixed with 5 µl of 18% PEG 6000, 0.1 M sodium citrate, pH 6.0 and equilibrated against 1 ml of 18% PEG 6000, 0.1 M sodium citrate, pH 6.0.

10 X-ray diffraction data was collected at 100K from native Ves v 5 crystals and after incorporation of heavy-atom derivatives and used to solve the three-dimensional structure of Ves v 5, see Figure 10 (manuscript in preparation).

Immunoelectrophoresis using rabbit polyclonal antibodies

The two Ves v 5 mutants were produced as recombinant Ves v 5 proteins and tested for their reactivity towards 20 polyclonal rabbit antibodies raised against recombinant Ves v 5. When analysed by rocket immunoelectrophoresis under native conditions, the rabbit antibodies were able to precipitate recombinant Ves v 5 as well as both mutants, indicating that the mutants have conserved α -25 carbon backbone tertiary structure.

Inhibition of specific serum IgE

The IgE-binding properties of Ves v 5 mutants were compared to recombinant Ves v 5 in a fluid-phase IgEinhibition assay using a pool of serum IgE derived from vesoid venom alleraic patients.

The inhibition assay was performed as described above using biotinylated recombinant $Ves\ v\ 5$ instead of $Bet\ v\ 1.$

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Ves v 5 Lys72Ala mutant

Lysine in position 72 show a high degree of solventexposure (70%) and is located in a molecular surface patch common for Vespula antigen 5. The relative orientation and high degree of solvent exposure argued for that lysine 72 can be replaced by an alanine residue without distortion of the α-carbon backbone tertiary structure. In addition, as none of the naturally occurring isoallergen sequences have alanine in position 72, the substitution of lysine with alanine gives rise to a non-naturally occurring Ves v 5 molecule.

IgE-binding properties of Ves v 5 Lys72Ala mutant

The IgE-binding properties of Ves v 5 Lvs72Ala mutant was compared with recombinant Ves v 5 in a fluid-phase IgEinhibition assay using the pool of serum IgE derived from birch allergic patients described above.

Figure 14 shows the inhibition of the binding of biotinylated recombinant Ves v 5 to serum IgE from a pool of allergic patients by non-biotinvlated Ves v 5 and by Ves v 5 Lys72Ala mutant.

There is a clear difference in the amount of respective recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant Ves v 5 reaches 50% inhibition at about 6 ng whereas the corresponding concentration for Ves v 5 Lys72Ala mutant is 40 ng. This show that the single point mutation introduced in Ves v 5 Lys72Ala mutant lowers the affinity for specific serum IqE by a factor of about 6. The maximum level of inhibition reached by the $Ves\ v$ 5 Lys72Ala mutant significantly lower

compared

recombinant Ves v 5. This may indicate that after the Lys72Ala substitution, some of the specific IgE present in the serum pool are unable to recognise the Ves v 5 Lys72Ala mutant.

Ves v 5 Tyr96Ala mutant

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Tyrosine in position 96 show a high degree of solvent-exposure (65%) and is located in a molecular surface 10 patch common for Vespula antigen 5. The relative orientation an high degree of solvent exposure argued for that tyrosine 96 can be replaced by an alanine residue without distortion of the three-dimensional structure. In addition, as none of the naturally occurring isoallergen 15 sequences have alanine in position 96, the substitution of tyrosine with alanine gives rise to a non-naturally occurring Ves v 5 molecule.

IgE-binding properties of Ves v 5 Tyr96Ala mutant

The IgE-binding properties of Ves v 5 Tyr96Ala mutant was compared with recombinant Ves v 5 in a fluid-phase IgE-inhibition assay using the pool of serum IgE derived from birch allergic patients described above.

Figure 14 shows the inhibition of the binding of biotinylated recombinant $Ves\ v$ 5 to serum IgE from a pool of allergic patients by non-biotinylated $Ves\ v$ 5 and by $Ves\ v$ 5 Tyr96Ala mutant.

There is a clear difference in the amount of respective recombinant proteins necessary to reach 50% inhibition of the binding to serum IgE present in the serum pool. Recombinant Ves v 5 reaches 50% inhibition at about 6 ng whereas the corresponding concentration for Ves v 5 Tyr96Ala mutant is 40 ng.

This show that the single point mutation introduced in $Ves\ v$ 5 Tyr96Ala mutant lowers the affinity for specific serum IgE by a factor of about 6.

The maximum level of inhibition reached by the Ves v 5
Tyr96Ala mutant significantly lower compared to
recombinant Ves v 5. This may indicate that after the
Tyr96Ala substitution, some of the specific IgE present
in the serum pool are unable to recognise the Ves v 5
Tyr96Ala mutant.

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CLAIMS

- 1. Recombinant allergen, characterised in that it is a non-naturally occurring mutant derived from a naturally occurring allergen, wherein at least one surface-exposed, conserved amino acid residue of a B cell epitope is substituted by another residue which does not occur in the same position in the amino acid sequence of any known homologous protein within the taxonomic order from which said naturally occurring allergen originates, said mutant allergen having essentially the same α -carbon backbone tertiary structure as said naturally occurring allergen, and the specific IgE binding to the mutated allergen being reduced as compared to the binding to said naturally occurring allergen.
- Recombinant allergen according to claim 1, characterised in that it is obtainable by
- a) identifying amino acid residues in a naturally occurring allergen which are conserved with more than 70% identity in all known homologous proteins within the taxonomic order from which said naturally occurring allergen originates;

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- b) defining at least one patch of conserved amino acid residues being coherently connected over at least 400 Å of the surface of the three-dimensional structure of the allergen molecule as defined by having a solvent accessibility of at least 20 %, said at least one patch comprising at least one B cell epitope; and
- c) substituting at least one amino acid residue in said at least one patch by another amino acid being non-conservative in the particular position while essentially preserving the overall α-carbon backbone tertiary

structure of the allergen molecule.

- Recombinant allergen according to claim 1 or 2, characterised in that the specific IgE binding to the mutated allergen is reduced by at least 5%, preferably at least 10%.
- 4. Recombinant allergen according to any of claims 1-3, characterised in that when comparing the α-carbon lobackbone tertiary structures of the mutant and the naturally occurring allergen molecules, the average root mean square deviation of the atomic coordinates is below 2A.
- 15 5. Recombinant allergen according to claim 2, characterised in that said at least one patch comprises atoms of 15-25 amino acid residues.
 - 6. Recombinant allergen according to any one of claims 2-5, characterised in that the amino acid residues of said at least one patch are ranked with respect to solvent accessibility, and one or more amino acids among the more solvent accessible ones are substituted.
- 5 7. Recombinant allergen according to claim 6, characterised in that one or more amino acid residues of said at least one patch having a solvent accessibility of 20-80 % are substituted.
- 8. Recombinant allergen according to any one of claims 2-7, characterised in that 1-5 amino acid residues per 400 Ų in said at least one patch are substituted.
- Recombinant allergen according to any one of claims
 2-5, characterised in that the substitution of one or more amino acid residues in said B cell epitope or said

- at least one patch is carried out by site-directed mutagenesis.
- 10. Recombinant allergen according to any one of claims 5 1-9, characterised in that it is derived from an inhalation allergen.
- 11. Recombinant allergen according to claim 10, characterised in that it is derived from a pollen 10 allergen.
 - 12. Recombinant allergen according to claim 10, characterised in that it is derived from a pollen allergen originating from the taxonomic order of Fagales, Oleales or Finales.
 - 13. Recombinant allergen according to claim 12, characterised in that it is derived from Bet v 1.
- 20 14. Recombinant allergen according to claim 13, characterised in that at least one amino acid residue of said B cell epitope or said at least one patch is substituted.
- 25 15. Recombinant allergen according to claim 14, characterised in that the substitution(s) is (are) Thr10Pro, Asp25Gly, (Asn28Thr + Lys32Gln), Glu45Ser, Asn47Ser, Lys55Asn, Thr77Ala, Pro108Gly or (Asn28Thr, Lys32Gln, Glu45Ser, Pro108Gly.

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- 16. Recombinant allergen according to claim 11, characterised in that it is derived from a pollen allergen originating from the taxonomic order of *Poales*.
- 35 17. Recombinant allergen according to claim 11, characterised in that it is derived from a pollen

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allergen originating from the taxonomic order of Asterales or Urticales.

- 18. Recombinant allergen according to claim 10, 5 characterised in that it is derived from a house dust mite allergen.
- 19. Recombinant allergen according to claim 18, characterised in that it is derived from a mite allergen originating from Dermatophagoides.
 - 20. Recombinant allergen according to claim 10, characterised in that it is derived from a cockroach allergen.
 - 21. Recombinant allergen according to claim 10, characterised in that it is derived from an animal allergen.
- 20 22. Recombinant allergen according to claim 21, characterised in that it is derived from an animal allergen originating from cat, dog or horse.
- 23. Recombinant allergen according to any one of claims 25 1-9, characterised in that it is derived from a venom allergen.
 - 24. Recombinant allergen according to claim 23, characterised in that it is derived from a venom allergen originating from the taxonomic order of Hymenoptera.
 - 25. Recombinant allergen according to claim 24, characterised in that is derived from a venom allergen from the taxonomic order of Vespidae, Apidae and Formicolide.

- 26. Recombinant allergen according to any one of claims 23-25, characterised in that it is derived from $Ves\ v\ 5$.
- 27. Recombinant allergen according to any one of claims 5 23-26, characterised in that at least one amino acid is substituted.
- Recombinant allergen according to any one of claims 25-27, characterised in that the substitution is Lys72Ala
 or Tyr96Ala.
 - 29. A method of preparing a recombinant allergen according to any one of claims 1-29, characterised by
- a) identifying amino acid residues in a naturally occurring allergen which are conserved with more than 70% identity in all known homologous proteins within the taxonomic order from which said naturally occurring allergen originates;

- b) defining at least one patch of conserved amino acid residues being coherently connected over at least 400 Å² of the surface of three-dimensional structure of the allergen molecule as defined by having a solvent 25 accessibility of at least 20%, said at least one patch comprising at least one B cell epitope; and
 - c) substituting at least one amino acid residue in said at least one patch by another amino acid being non-conservative in the particular position while essentially preserving the overall α -carbon backbone tertiary structure of the allergen molecule.
- 30. A method according to claim 29, characterised by ranking the amino acid residues of said at least one patch with respect to solvent accessibility and

substituting one or more amino acids among the more solvent accessible ones.

- 31. A method according to claim 29 or 30, characterised 5 in that the substitution of one or more amino acid residues in said B cell epitope or said at least one patch is carried out by site-directed mutagenesis.
- 32. Recombinant allergen according to any of claims 1-28 for use as a pharmaceutical.
- 33. Pharmaceutical composition, characterised in that it comprises a recombinant allergen according to any one of claims 1-28, optionally in combination with a pharmaceutically acceptable carrier and/or excipient, and optionally an adjuvant.
 - 34. A pharmaceutical composition according to claim 33, characterised in that it is in the form of a vaccine against allergic reactions elicited by a naturally occurring allergen in patients suffering from allergy.
- 35. Method of generating an immune response in a subject comprising administering to the subject at least one recombinant allergen according to any one of claims 1-28 or a pharmaceutical composition according to any one of claims 33-34.
- 36. Process for preparing a pharmaceutical composition according to any one of claims 33-34 comprising mixing at least one recombinant allergen according to any one of claims 1-28 with pharmaceutically acceptable substances and/or excipients.
- 35 37. Vaccination or treatment of a subject comprising administering to the subject at least one recombinant

allergen according to any one of claims 1-28 or a pharmaceutical composition according to any one of claims 33-34.

5 38. Pharmaceutical composition obtainable by the process according to claim 36.

39. Method for the treatment, prevention or alleviation of allergic reactions comprising administering to a subject a recombinant allergen according to any one of claims 1-28 or a pharmaceutical composition according to any one of claims 33-34 or 38.

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Fig. 1

non-sense primer	3'-	ATACTCTGACTCTGGGGGAGACA	-5'
sense primer	5'-	TGAGACCCCCTCTGTTATCCCAG	-3 '
Bet v 1 non-sense	3'-	${\tt TTAATACTCTGACTCTGG\underline{T}GGAGACAATAGGGTCGTCGAGC}$	-5'
Bet v 1 sense	5'-	AATTATGAGACTGAGACCACCTCTGTTATCCCAGCAGCTCG	-3'

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Fig. 2

Oligonucleotide primers for site directed mutagenesis of Bet v 1 (No. 2801).

all	sense	1: 183Bv, 15-mer 5'-GTTGCCAACGATCAG
1	sense	2: 184Bv, 23-mer 5'-TGAGACCCCCTCTGTTATCCCAG
1	non-sense	3: 185Bv, 23-mer 5'-ACAGAGGGGGTCTCAGTCTCATA
2	sense	4:1868v, 31-mer 5'-GATACCCTCTTTCCACAGGTTGCACCCCAAG
2	non-sense	5: 187Bv, 31-mer 5'-ACCTGTGGAAAGAGGGTATCGCCATCAAGG
3	sense	6: 188Bv, 23-mer 5'-AACATTTCAGGAAATGGAGGGCC
3	non-sense	7: 189Bv, 23-mer 5'-TTTCCTGAAATGTTTTCAACACT
4	sense	8: 190Bv, 23-mer 5'-TTAAGAACATCAGCTTTCCCGAA
4	non-sense	9: 191Bv, 23-mer 5'-AGCTGATGTTCTTAATGGTTCCA
5	sense	10: 192Bv, 23-mer 5'-GGACCATGCAAACTTCAAATACA
5	non-sense	11: 193Bv, 23-mer 5'-AGTTTGCATGGTCCACCTCATCA
6	sense	12: 194Bv, 23-mer 5'-TTTCCCTCAGGCCTCCCTTTCAA
6	non-sense	13: 195Bv, 23-mer 5'-AGGCCTGAGGGAAAGCTGATCTT
7	sense	14: 196Bv, 24-mer 5'-TGAAGGATCTGGAGGGCCTGGAAC
7	non-sense	15: 1978v, 24-mer 5'-CCCTCCAGATCCTTCAATGTTTTC
8	sense	16: 198Bv, 24-mer 5'-GGCAACTGGTGATGGAGGATCCAT
8	non-sense	17: 199Bv, 24-mer 5'-CCATCACCAGTTGCCACTATCTTT
all	non-sense	18: 200Bv, 15-mer 5'-CATGCCATCCGTAAG

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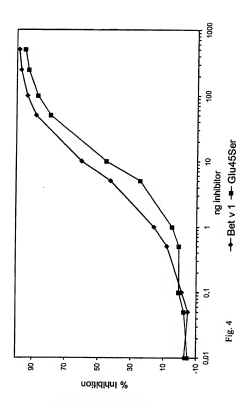
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Fig. 3

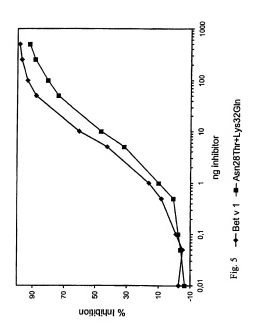
Overview of all Bet v 1 mutations

1(A-C) GGTGTGTTTAATTATGAGACTGAGACCACCTCTGTTATCCCAGCAGCTCGACTGTTCAAG 60 G V F N Y E T E TT-PS V I P A A R L F K 20 9(A-G) 2(A-C) 2(A-C) GCCTTTATCCTTGATGGCGATAACCTCTTTCCAAAGGTTGCACCCCAAGCCATTAGCAGT 120 A F I LD-GG DN-TL F PK-QV A P Q A I S S 40 4 (G-C) 6 (GA-TC) 3 (GA-TC) 7 (AA-TC) GTTGAAAACATTGAAGGAAATGGAGGGCCTGGAACCATTAAGAAGATCAGCTTTCCCGAA V E N I E-S G N-S G G P G T I K K-N I S F P E-S 5 (CA-TG) GGCCTCCCTTTCAAGTACGTGAAGGACAGAGTTGATGAGGTGGACCACACAAACTTCAAA 240 G L P F K Y V K D R V D E V D H T-A N F K 80 TACAATTACAGCGTGATCGAGGGGGGTCCCCATAGGCGACACATTGGAGAAGATCTCCAAC 300 YNYSVIEGGPIGDTLEKISN 100 10 (GAG-CAC) 8 (CCC-TGG) GAGATAAAGATAGTGGCAAC<u>CCC</u>TGATGGAGGATCCATCYTGAAGATCAGCAACAAGTAC 360 EIKIVATP-GDGGSILKISNKY CACACCAAAGGTGACCATGAGGTGAAGGCAGGCAGGTTAAGGCAAGTAAAGAAATGGGC 420 H T K G D H E V K A E Q V K A S K E M G 140 GAGACACTTTTGAGGGCCGTTGAGAGCTACCTCTTGGCACACTCCGATGCCTACAACTAA 480

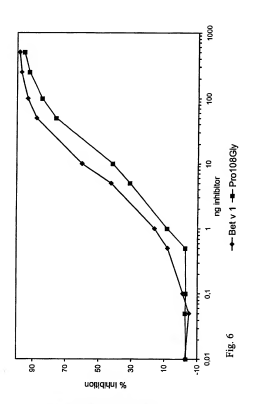
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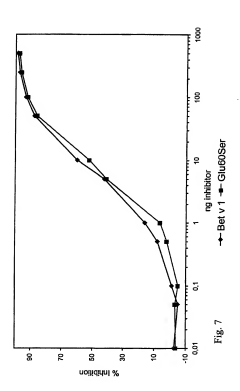
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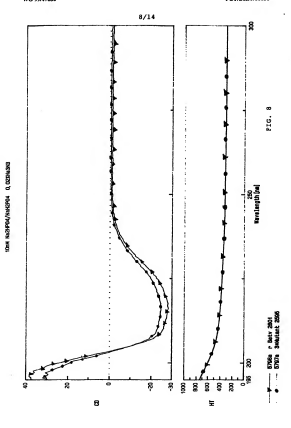
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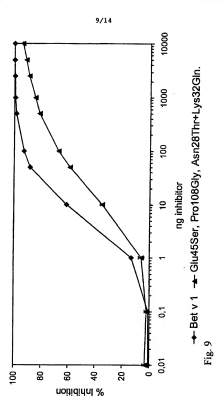
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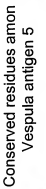
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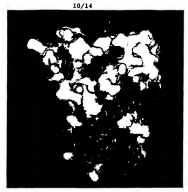
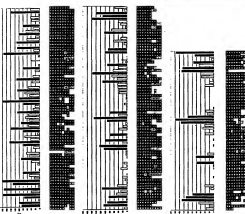


Figure 10.



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Mutant-specific oligonucleotide primers used for Ves v 5 mutants. Mutated nucleotides underlined.

Ves v 5 mutant 1 (K72A)

Ves v 5 sense 5" - ACCACACCTCCAGCAGAGATATGAAAATTTOGTATGGA -3 Ves v 5 non-sense 3" - TGGTGTCGGAGGTGGCTTCTTATACTTTTTAACCATACCT -5 sense primer 5" - CCAGGGGTATATGAAAAT -3 non-sense primer 3" - GTCGGAGGTGGCGGATTATAC -5			_
Ves v 5 non-sense 3'- TGGTGTCGGAGGTCGCTTCTTATACTTTTTAAACCATACCT -5	non-sense primer	3'- GTCGGAGGTCGC <u>CGA</u> TTATAC -	51
	sense primer	5 - CCAGCG <u>GCT</u> AATATGAAAAAT -:	3 -
Ves v 5 sense 5'- ACCACAGCCTCCAGCGAAGAATATGAAAAATTTGGTATGGA -3	Ves v 5 non-sense	3 - TGGTGTCGGAGGTCGCTTCTTATACTTTTTAAACCATACCT -	5-
	Ves v 5 sense	5 - ACCACAGCCTCCAGCGAAGAATATGAAAAATTTGGTATGGA -	3 -

Ves v 5 mutant 2 (Y96A)

ves	٧	9	sense		GGCIANICANIGICANIAIGGICACGAIACIIGCAGGGAIG	-3
Ves	v	5	non-sense	3	CCGATTAGTTACAGTTATACCAGTGCTATGAACGTCCCTAC	-5
sens	3e	p	rimer	5′-	TGTCAAGCTGGTCACGATACT	-31
non-	- 56	ens	se primer	3^-	TTAGTTACAGTT <u>CG</u> ACCAGTG	-51

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Fig. 12

Oligonucleotide primers for site directed mutagenesis of $Ves\ v\ 5.$

all sense 1: XhoI start, 38-mer:

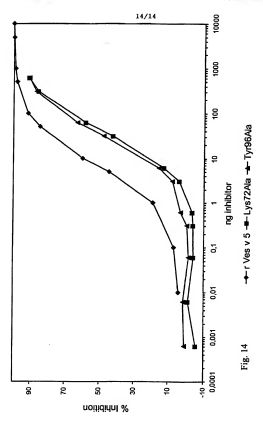
BCORI 5'-CCGCTCGAGAAAAGAAACAATTATTGTAAAATAAAATG L R R N Y C K I K . Kex2 cleavage site amino terminus of Ves v 5

1	sense	1: K7	2As 21-	mer	5 - CCAGCGGCTAATATGAAAAAT
1	non-sense	2: K7	2Aa 21-	mer	5 ~- CATATTAGCCGCTGGAGGCTG
2	sense	3: Y9	6As 21-	mer	5 -TGTCAAGCTGGTCACGATACT
2	non-sense	4: Y9	6Aa 21-	mer	5 - GTGACCAGCTTGACATTGATT
all	non-sense	7: CT	-nPTCZaA	21-mer	5 ~ ATTCATCAGCTGCGAGATAGG

Fig.	13	13/14

Outomrieu	٥f	Vee	7.	_	mutations

1	AA	CAA	TTA	TTG	TAA	AAT	AAA	ATG	TTT	GAA	AGG	AGG'	TGT	CA	FAC	TGC	CTG	CAA	TA	rgga	60
1	N	N	¥	С	K	1	K	c	L	K	G	G	v	н	T		C		Y		20
																			Ī	•	
61	AG	CT	raa.	ACO	GAA:	rrg	CGG	TAA	TAA	GT.	AGTO	GT	ATC	TA	rgg:	ст	AACC	IAA.	CAI	AGAG	120
21	. s	L	K	P	N	С	G	N	ĸ	ν	v	ν	s	Y	G	L	т				40
													-	-	-	_	•	•	۳	_	
121	AA	CA	AGA	CAT	CTT	AA	GGA	3CA	CAA	rgac	TT	rag)	ACA?	LAA.	ATT	rgcz	\CG/	GGJ	TTC	GAG	180
41	K	Q	D	1	L	K	E	H	N	D	F	R	Q	K	1	A	R	G	L	E	60
														A]							
181	AC.	AG	\GG'	TAA:	rcc1	CGG	ACCI	CAC	GCC7	rccz	AGCC	AAG	AAT	ATO	AA.	LAAT	TTC	GTA	TGC	AAC	240
61	T	R	G	N	P	G	P	Q	P	P	A	ĸ	N	М	K	N	L	V	W	N	80
																21	T96	AT 6	TA-	GC)	
241	GAC	GAC	TT	AGC	TAT	GTO	GCC	CA	GTC	TGG	GCI	CAA:	CAA	TGT	CAA	TAT	GGT	CAC	GAT	ACT	300
81	D	Е	L	A	Y	v	А	0	v	W	A	N	0	c	0	¥	G	н	D	т	100
								_					-		-		_		-	•	100
301	TGC	AGG	GA?	GT	.GCA	AAZ	TAT	CAC	GTT	GGA	CAA	AAC	GTA	GCC	тта	ACA	сст	ממר	a Cro	GCT	360
101	С	R	D	v	А	K	Y	0	ν	G	0	N	v	A	T.	т	G			Δ.	120
								-			•				_	-	•	·	•	•	120
361	GCT	AAA	TAC	GAT	GAT	CCF	GTI	AAA	CTA	GTI	AAA	ATG	TGG	GAA	GAT	GAA	GTG	AAA	CAT	TAT	420
121	A	ĸ	Y	D	D	P	v	K	L	v	к	м	W	R			v				140
														_	-	_			-	•	
421	AAT	сст	AAG	AAA	AAG	TTI	TCG	GGA	AAC	GAC	TTT	CTG	AAA	ACC	GGC	CAT	TAC	ACT	CAA	ATG	480
141	N	P	K	ĸ	K	F					F						Y		0		160
																			-		
481	GTT	TGG	GCT	AAC	ACC	AAG	GAA	GTT	GGT	TGT	GGA	AGT	ATA.	AAA	TAC	ATT	CAA	GAG	AAA	TGG	540
161	٧	W	A	N	T	ĸ	Е	ν	G	C	G	s	I	ĸ	Y	I	0	E	ĸ	W	180
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541	CAC	AAA	CAT	TAC	CTT	GTA	TGT	AAT	TAT	GGA	ccc	AGO	GGA	AAC	TT.	AAG	AAT	GAG	SAA	TT	600
181	H	K	H	Y	L	٧	C	N	Y	G	P	s	G	N	F	K	N	B	E	L	200
601																					612
201	Y	O	т	ĸ	ati	nn															204



SUBSTITUTE SHEET (RULE 26)

INTERNATIONAL SEARCH REPORT

Int. _at Application No

PCT/DK 99/00136 A. CLASSIFICATION OF SUBJECT MATTER
TPC 6 C12N15/29 C12N15/12 CO7K14/415 CO7K14/435 A61K39/35 A61K39/36 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED (classification system followed by classification symbols) IPC 6 C12N CO7K Documentation searched other than minimum documentation to the axient that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category * Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X FERREIRA F ET AL.: "Modulation of IgE 1-14,16, reactivity of allergens by site-directed 29-39 mutagenesis: potential use of hypoallergenic variants for immunotherapy" FASEB JOURNAL FOR EXPERIMENTAL BIOLOGY, vol. 12, no. 2, February 1998 (1998-02), pages 231-242, XP002085249 BETHESDA, MD US cited in the application page 240, right-hand column; figures 1.3 -/--X Further documents are lated in the continuation of box C. X Patent family members ere listed in annex. isi categories of cited documents : later document published after the international fit or priority date and not in conflict with the applicated to understand the principle or theory under trivention. "A" document defining the general state of the art which is not considered to be of particular relevance earlier document but published on or after the international filing date document of particular reliocument which may throw doubts on priority claim(e) or which is clied to establish the publication date of another citation or other special reason (as specified) document referring to an oral disclosure, use, exhit other means document published prior to the international fiting date but later than the priority date claimed "&" document member of the same patent territy Date of the actual completion of the international search Date of mailing of the international search report 30/08/1999 24 August 1999 no address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentisan 2 Nt. - 2280 HV Rijewijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nt. Fax: (+31-70) 340-3016 Cupido, M

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bs. sel Application No PCT/DK 99/00136

	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
		PROPERTY OF CHIEF INC.
'	WIEDEMANN P ET AL.: "Molecular and structural analysis of a continuous birch profilin epitope defined by a monoclonal antibody"	1-12, 29-39
	JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 271, no. 47, 22 November 1996 (1996-11-22), pages 29915-29921, XP002085250	
ĺ	MD US cited in the application page 29919; figure 6; table I	
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	l November 1993 (1993-11-01), pages 707-716, XP002035181 ISSN: 0091-6749 page 715, left-hand column	

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INTERNATIONAL SEARCH REPORT

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PCT/DK 99/00136

Box (Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)
This int	ernational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
t. X	Claims Nos: because thy relate to subject matter not required to be searched by this Authority, namely: Remark: Although claims 35, 37 and 39 are directed to a method of treatment of the human or animal body, the search has been carried out and based on the alleged effects of the composition.
2.	Claims Nov. Better to parts of the International Application that do not comply with the prescribed requirements to such because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, specifically.
з 🗌	Claims Not.: . Leanuse they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a),
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
t	As all required additional search fees were timely paid by the applicant, this international Search Report covers all searchable claims.
2. 🗌	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search lies were timely paid by the applicant, this international Search Report covers only those claims for which fiers were paid, specifically claims Nos;
4.	No required additional search frees were timely paid by the applicant. Consequency, this international Search Report is restricted to the invention first inventioned in the clasmic, it is convered by claims. Noc.:
Remark	on Protest The additional search tres were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT Information on patent family members PCT/DK 99/00136												
Patent document cited in search repor	t	Publication date	P	atent family member(s)	Ý	Publication date						
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